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Glykosid ester and their production as well as use in cosmetics,

Pharmaceutical products and Nahrungs-bzw. Feeds

The present invention concerns new, biological active < RTI ID=1.1> Glykosid ester, < /RTI> Method to their production, these junctions contained cosmetic and/or pharmaceutical preparing as well as these bonds contained Nahrungs-und feed.

In the Kosmetik becomes ever more important the application of active substances. With the active substances, which so far already find application in the Kosmetik, it does not concern always natural substances. The optimization of known active substances and the production of new active substances are subiect matter of many research work.

In the broadest sense active substances such cloths are occurring, those in relatively small quantities or supply-large physiological effect to unfold can. Here is to hormones, vitamins, enzymes, trace elements etc. to think, in addition, of medicines (drugs), < RTI ID=1.2> Fodder additives, Düngemittels / RTI> and pesticides. Pretty often one can observe also svererism.

Glykoside and < RTI ID=1.3> arvlaliphatische Glykosidester</RTI> Those inventive accomplished derivatizations obtain an improved effect as well as an increased bioavailability, how it became already early shown at the example of Salicinderivate.

Many natural occurring Alkyl-und of phenol glucosides show antiviral, antimicrobial and partial antiinflammatorische effects. They are however often bioavailable due to their polarity a little and/or, their selectivity is too small.

For example Salicin (a plycosidic active substance from the pasture crust) is < RTI ID=1.4> nichtsteroidales antinifammatorischeex./RTI > agent (NSAIA), which shows remarkably improved effectiveness after derivatization (esterifications). Recently the synthesis new aryl-aliphatic Salicinester succeeded such as Phenylacetoy) Salicin, whereby the esterification preferred at the primary OH-groups of the Salicinic (first at the surger than at the benzyle remainder) in the Salicin made. Due to the aryl-aliphatic remainder the material transfer is improved that be supported to the surger of the surger o

In addition Ericaceae became, z from excerpts of the plants of the species. B. from the bear grape/cluster (Arctostaphylos wu arei L.). The Glykopid Arbutin insulated, which strike-lightening effect shows. This Glykopid < a certain: RTI ID=2.5 biotechnologischese: /RTI> Interest is used, there it in Japan because of its lightening effect under the Inhibiterrung of the Metaninbiosynthese (inhibition of the key enzyme tyrosianse) in considerable quantity (1-2 t per year) in cosmetics. While in the European market the preparations rather approximately < RTI ID=2.6 > Old marks, liver flockener / RTI> and/or.

& top Summer rungs used are <, are in the asiatic market the ideals of beauty of one; RTI ID=2.7> bright makellosen< /RTI> Skin by whole-physical treatment achieved become.

« RTI ID-2.8» Alkvlpohydµuoside« (RTI» (APG) Alkylpohygluooside (APG), become as phosphate-free Neutraltenside and. A. Detergents and cosmetics added (on the basis of regenerating raw materials). 1-7 glucose units is glycosidic linked with a Fettalkohol (mostly 12 C-atoms):

EMI2.1

PUFAs and CLAs natural one of oils (z. B. from sunflower oil, linseed oil, c RTI ID-2.9- Oil tree oil) x (RTI), with an high proportion at polyungestitigen latty acids (English: polyunsaturated fatty acids, PUFAs) become used in the Kosmetik and Dermatologie. The PUFAs belongs in the diet to the group of the essential fatty acids and shows additionally a positive effect with the tray in the prophylaxis of arteriosclorosis. Sesides are also pharmaceutical effects of importance: They can do antiinflammatorische (inhibition Prostaglandin-bzw. Leukotriensynthese), in addition, a thrombolvitische and hypotensische effect achtibit.

Inventive one becomes PUFA defined as a multiple installated fatty acid with 16 to 26 -atoms, whereby the fatty acid exhibits at least four insulated and/or at least wo onjugated double bonds. Examples for PUFAs are altogether twelve < to the Linelsäure those; RTI ID-3.15 (cis., </RT> (cis., </rr> (cis., </rd>

 composition most different to CLA mixtures in dependence of the reaction conditions (z. B. Edenor UKD 6010, < RTI ID=3.2> Handle KGaA), < /RTI>

Due to its conjugated double bonds this isomers Octadecadiensäuren becomes also as " conjugated linoleic acids " (CLAs) referred.

One receives it to natural way as mixture major from milk and meat from ruminants (5 mg/g < RTI ID=3.3> Fat), < /RTI> in those it by the Pansenbakterium Butyrivibrio < RTI ID=3.4> fibrisolvens< /RTI> are synthesized.

The effect of the CLAs is very versatile. They show inhibitive effect with the Karzinogenese and with the Artherogenese, bow further an anti-oxidative effect by Furanbildung, which plays an important role with the prevention of crayfish and koronaren heart diseases (KHKen). The Vermehrung Muskel-und bone mass through < RTII D=3.5> Fetdeport reduzierende

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Fatty acid oxidizing in Muskel-und < RTI ID=4.2> Fettzellen< /RTI> recycled become.

A further application of these CLAs than isomeric mixture follows the precursor of the Entzündungsmediators from their competent effect than antiinflammatorische agents, there it due to their structural distorting shank to the arachidonic

Prostaglandin, die zur Biosynthese der Prostaglandine benötigten Desaturasen kompetitiv hemmen können. They displace additionally the precursors Arachidon and

Eicosansäure aus den Phospholipiden und hemmen die Umwandlung der Arachidonsäure in die Eicosansäure und damit die Umwandlung in die Prostaglandine.

Although in the literature already numerous pharmakologisch effective cloths are described, those for example in < RTI | D=4.3> Entzündungskaskade< /RTI> engage, exists turther a need at better effective active substances poor in side effects.

A need at active substances with a good absorbableness and a quick penetration continues to exist into the skin, which besides good into pharmaceutical or cosmetic formulations be trainable must.

The object, which is the basis for the present invention, consisted thus of it, such side effect-poor, good working and good those which can be processed and too < RTI ID=4.4> applizierenden
 /RTI> To make available materials.

RTI ID-4.55 Glykosides / RTI> and certain
RTI ID-4.65 Glykosidester / RTI> are z. B. aus den Natur bekannt.
Known (neither from plants, microorganisms or animal cells sith synthetic prepared) are not however the here described seters, in particular PUFA and CLA ester, from insatiated fatty acids with sugars or Glykosiden, with those at least one of the hydroxyl groups of the sugar with (insatiated) Carbonbzw. Fatty acid third in State of the control of the control

Überraschenderweise wurde von den Erlindern gelunden, dass bestimmte Ester, insbesondere Ester von ungseättigten Fettsbauren mit Zuckern der Glykosiden eine gegenüber den bekannten Einzelkomponenten (Fettsbaure bzw. « RIT ID-4.7 Sugar/Glykosid) « /RIT» improved biological availability, amplitied effect and/or a broadened effect spectrum exhibt. Peleraby made « RIT ID-4.8 Se teströllung» / RIT» einttels der primären Hydroxylgruppe des Zuckers/Glykosids; however also the remaining OH-groups at the sugar or eventual existing OH-groups in « can; RIT ID-5.15. Algkote.) kMTov (z.e. Netroverset in the case of the Arbutins) for the esterification to be used.

By the provision of the bonds of the present invention the inventors knew solve the problem posed.

The connections of the present invention are esters < RTI ID=5.2> alignmeninen formula (1) < /RTI> : < RTI ID=5.3> ACA CO 2 or ". (1), < /RTI> > where R a hydrogen, a one < over one; RTI ID=5.4> Etherbrückee /RTI> at the sugar bound branched or straight-chain < RTI ID=5.5> CB-C20-AB/yl, < /RTI> one Aylalkyl or a substituted or a unsubstituted or a lord straight-chain < RTI ID=5.5> CB-C20-AB/yl, < /RTI> one Aylalkyl or a substituted or a disaccharide or a polysaccharide, which are n-multiple ester-like with AC substituted and, if R does not stand for hydrogen, acetalisch to the remainder, Bhotid, where AC a multiple insaltated < RTI ID=5.7> CF-C25-Acytrest <RTI> with AC substituted and/ or at least two conjugated double bonds or it represents an aryl-aliphatic remainder with 1-q groups of groupproups between group of esters and aromatic ring where m an integral number (12.3-7), including 0 if its where n an integral number (12.3-7), bowever 0, is not <, under the condition; RTI ID=5.8> that Z-O-Rmc /RTI> not for Saldicin, which is <RTI ID=5.9> Liftydrownerthy-phenyl) - sex /RTI 3 glucopyranosid, steht

Bevorzugt als Fettsäuren sind CIAs und Stearidonsäure, besonders bevorzugt die Octadecadiensäuren, die über konjugierte Doppeblindungen an den C-Atomen 9 und 11,10 und 12 oder 11 und 13 verfügen, insbesondere die cis, trans, 9,11-und die cis, cis, 9,11-Octadecadiensäure. EMIS 1

Stearidonsāure (CAS 20290-75-9); 6.9.12.15-Octadecatetraenoic acid, (6Z, 9Z, < RTI ID=5.10> 1 2Z, 1 5Z) - (9CI) < /RTI>

As sugar mono, Di-und Oligosaccharide, in particular D glucose, come D

Anteil (Z-O-Rm) Fragilin</RTI> (d. h. m = 1). AC the acvl radical cis. trans.

Galactose, D-xylose, D-Apiose, L-Rhamnose, L-arabinose and Rutinose, in considerations, whereby D glucose is particularly prefered.

Als Glykoside (Z-C-Rm) kommen als Zuckerbestandteil Monosaccharide (wie die im vorhergehenden Absatz genannten) enthaltende Verbindungen, insbesondere Arbutin, <RTI ID=6.1> Fragilin</RTI> und Poplin, aber auch Oligo-oder Polysaccharide enthaltende Glykoside wie

Alkylpoly glucoside, particular APG (handle < RTI ID=6.2> KGaA), < /RTI> in considerations.

Geeignet im Sinne der Erfindung sind Verbindungen, bei denen der Glykosid-Anteil (Z < RTI I D=6.3>O-Rm)</RTI>
Arbutin (D. h. m = 1), AC the acyl radical cis, trans, 9,11-oder cis, cis, 9,11
Octadecadiensäure and n 1 are, Gleichfalls geeignet sind solche Verbindungen, bei denen der < RTI I D=6.4>Glykosid-

9,11-oder cis, cis, 9,11-Octadecadiensaure and n of the 1 is.

Other favourable bonds are esters, whereby the Glykosid portion < RTI ID=6.5> (Z-O-Rm) < /RTI> Arbutin (D. h. m = 1), Ac der Acylrest der cis, trans, 9,11-oder der cis, cis, 9,11 Octadecadiensäure und n 2 ist; Ester, whereby < RTI ID=6.6> Glykosid portion (Z-O-Rm) Fragilin< /RTI> (d. h. m = 1), AC the acyl radical cis, = trans, 9.11-oder cis, cis, 9.11-Octadecadiensäure and n of the 2 is: Verbindungen, bei denen der Glykosid-Anteil < RTI ID=6.7 > (Z-O-Rm) < /RTI > Arbutin or Fragilin (D. h. m = 1). AC the acyl radical of the Stearidonsaure and n 1 is; schliesslich Verbindungen, wobei der Glykosid-Anteil < RTI ID=6.8>(Z-O-Rm)</RTI> Arbutin or Fragilin (D. h. m = 1), Ac der Acylrest der Stearidonsäure und n 2 ist.

Other favourable bonds are esters with the subsequent parameters; if n = 1 is, the single remainder of AC to a primary OH-group of the sugar bound, in particular to that primary OH-group, which sits at the sugar molety, is bound to which also B, if m < RTI ID=6.9> w< /RTI> 0; if n = 2 is, a remainder of AC to that primary OH-group, which sits at the sugar moiety, is bound to which also B, if m < BTI ID=6.10 > X< /BTI> 0, and the other remainder of AC is to a further primary or to a secondary OH-group of the sugar bound.

Particularly prefered bonds according to the present invention are the subsequent esters 1 to 30, with which both m and n are 1, with those AC, Z and R the subsequent importances have and with those the ester group over the primary alcohol group of the sugar/Glykosids formed become (are several Glucose-bzw.

Zucker-Einheiten vorhanden, liegt die Veresterung am primären OH der Glucose Einheit vor, die auch glykosidisch verknüpft ist mit z. B. <RTI ID=7.1>CH3-(CH2) 10-CH2OH)</RTI>

- 1. AC is cis, trans, 9,11-Octadecadienoyl; Z is D glucose; is H; 2. AC is cis, cis, 9,11-Octadecadienoyl; Z is D glucose; R is H:
- 3. AC is trans, cis, 10,12-Octadecadienoyl; Z is D glucose; R is H; 4. AC is cis, trans, 9,11-Octadecadienoyl; Z is D glucose; Rist-para-C6H4-OH;
- 5. AC is cis, cis, 9.11-Octadecadienoyl; Z is D glucose; R ist-para-C6H4-OH; 6. AC is trans, cis, 10,12-Octadecadienoyl; Z is D glucose; R < RTI ID=7.2> ist-para-C6H4-OH< /RTI>; 7. Ac ist cis, trans, 9.11-Octadecadienoyl; Z is D glucose;
- R is < (CH2): RTI ID=7.3> rxCH3< /RTI>
- 8. AC is cis. cis. 9.11-Octadecadienovl; Z ist D-Glucose; R ist (CH2) < RTI ID=7.4>11CH3</RTI> 9. Ac ist trans. cis. 10.12-OctadecadienovI: Z is D glucose: R ist < RTI ID=7.5> (CH2) < /RTI> < RTI ID=7.6>
- 11CH3< /RTI>
- 10. AC is cis. trans. 9.11-Octadecadienovl: Z ist (D-Glucose) 2: R ist (CH2) < RTI ID=7.7>1rCH3</RTI>: < RTI ID=7.8> 11.< /RTI> AC is cis, cis, 9.11-Octadecadienoyl; Z is < (D glucose); RTI ID=7.9> 2< /RTI>; R is < (CH2);
- RTI |D=7.10> arCH3< /RTI> ; 12. AC is trans, cis, 10,12-Octadecadienoyl; Z is (D glucose) 2; R is < RTI ID=7.11> (CH2) r1CH3< /RTI>;
- 13. Ac ist cis. trans. 9.11-Octadecadienoyl; Z is (D glucose) 3; R ist (CH2) < RTI ID=7.12>11CH3</RTI>;
- 14. AC is cis, cis, 9,11-Octadecadienoyl; Z is (D glucose) 3; R ist (CH2) < RTI ID=7.13>11CH3</RTI>
- 15. AC is trans. cis. 10.12-Octadecadienovi; Z is (D glucose) 3; R is < (CH2); RTI ID=7.14> 11CH3< /RTI>;
- 16. Ac ist cis, trans, 9,11-Octadecadienoyl; Z is (D glucose) 4; R is < (CH2); RTI ID=7.15> 11CH3< /RTI >;
- 17. AC is cis, cis, 9,11-Octadecadienoyl; Z is (D glucose) 4; R is < (CH2); RTI ID=7.16> 11CH3< /RTI>
- 18. AC is trans, cis, 10,12-Octadecadienoyl; Z is (D glucose) 4; R is < (CH2); RTI ID=7.17> 11CH3< /RTI>
- 19. AC is cis, trans, 9,11-Octadecadienoyl; Z is (D glucose) 5; R is < (CH2); RTI ID=7.18> 11CH3< /RTI>; 20. AC is
- cis, cis, 9,11-Octadecadienoyl; Z is (D glucose) 5; R is < (CH2); RTI ID=7.19> 11CH3< /RTI>; 21. AC is trans, cis, 10.12-Octadecadienoyl; Z is (D glucose) 5; R is < RTI ID=7.20> (CH2) < /RTI> < RTI ID=7.21> 11CH3< /RTI> ; 22.
- AC is cis. trans. 9.11-Octadecadiencyl; Z is (D glucose) 6; R is < (CH2); RTI ID=7.22> 11CH3< /RTI>; 23. AC is cis. cis, 9,11-Octadecadienoyl; Z is (D glucose) 6; R ist (CH2) < RTI ID=7.23>11CH3</RTI>; 24. AC is trans, cis, 10,12-Octadecadienoyl; Z is (D glucose) 6; R is < (CH2); RTI ID=7.24> 11CH3< /RTI>; 25. AC is cis, trans, 9,11-
- Octadecadienoyl; Z is < RTI ID=7.25> (D glucose) 7< /RTI> ; R is < RTI ID=7.26> (CH2) nCH3< /RTI> ; 26. Ac ist cis, cis, 9,11-Octadecadienoyl; Z is (D glucose) 7; R is < (CH2); RTI ID=7.27> 11CH3< /RTI>; 27. AC is trans, cis, 10,12-Octadecadienoyl; Z is (D glucose) 7; R is < RTI ID=7.28> (CH2) < /RTI> < RTI ID=7.29> 11CH3< /RTI>; 28. AC is 6,9,12,15-Octadecatetraenoyl (6Z, 9Z, 12Z, 15Z); Z is D glucose; R is H; 29. AC is 6,9,12,15-Octadecatetraenoyl (6Z, 9Z, 12Z, 15Z); Z is D glucose; R be p < RTI ID=7.30> C6H4-OH< /RTI> ;
- 30. AC is 6.9.12.15-Octadecatetraenoyl (6Z, 9Z, 12Z, 15Z); Z is D glucose; and R is < RTI ID=8.1> (CH2) 11 CH3-< /RTI>
- Further prefered bonds are the esters 31 to 60, itself from the esters the 1 to < RTI ID=8.2> 30< /RTI> lediglich dadurch unterscheiden, dass die Zuckereinheit (en) nicht aus D-Glucose, sondern aus der entsprechenden Anzahl von D-Galactose-Einheiten besteht. The ester
- 54 is thus, in order to give an example, an analogue to the esters 24 and thus an ester, with the AC of trans, cis, 10,12-Octadecadienoyl; Z (D-Galactose) 6; R < RTI ID=8.3> (CH2) ?CH3< /RTI> ; m and n 1 are.

Prefered ones are also the subsequent esters 61 to 120 in analogy to the esters 1 to 60, which differ however from the latters by the fact that n = 2 and that the second AC remainder at the 4-OH Glucose-bzw. Galactose-Einheit sitzt.

Weiterhin bevorzugt sind die folgenden Ester 121 bis 180 in Analogie zu den Estern 61 bis 120, die sich allerdings von letzteren dadurch unterscheiden, dass der zweite Ac Rest nicht am 4-OH, sondern am 1-OH der Glucose-bzw. Galactose unit sits.

- Likewise prefered is the subsequent esters 181 to 240 in analogy to the esters
- 121 to 180, which differ however from the latters by the fact that the second AC remainder not at the 1-OH, but at the 2-OH Glucose-bzw, Galactose unit sits.

Weitere bevorzugte Ester sind die folgenden Ester 241 bis 300 in Analogie zu den Estern 1 bis 60, die sich allerdings von letzteren dadurch unterscheiden, dass n = 3 und dass diese < RTI ID=8.4> Verknüp-fungen</RTI> at the primary (6-OH) as well as the 1-OH-und of the 4-OH group of the sugar take place.

Other prefered esters are the subsequent esters 301 to 360 in analogy to the esters 241 to 300, which differ however from the latters by the fact that the three AC remainders at the primary as well as the 2-OH-und of the 4-OH-Gruppe of the sugar are bound.

Wiederum andere bevorzugte Ester sind die folgenden Ester 361 bis 420 in Analogie zu den Estern 1 bis 60, die sich allerdings von letzteren dadurch unterscheiden, dass n = 4 und dass diese Verknüptungen an der primären (6-OH-) sowie der 1-OH-, der 2-Ohund der 4-OH-Gruppe des Zuckers erfolgen.

By the inventive modification of the fatty acids, D. h. by the described above derivatization of the fatty acids in form of the junctions of the present invention, the compatibility as well as the biological availability and effect of such fatty acids for the inset in Kosmetik, pharmacy and/or diet are improved.

Dementsprechend betreffen weitere Aspekte der vorliegenden Erindung die Verwendung der Verbindungen der allgemeinen Formel «FIII ID» 3.1 (1)-«FIII > zur Herstellung von kosmetischen und pharmazeutischen Zubereitungen, ihre Verwendung als Additive zu Nahrungs- «FII ID» 9.2 × Nahrungsergänzungs- und«/RII» Feeds. Weitere Aspekte der vorliegenden Erifindung betreffen pharmazeutische und kosmetische Zubereitungen sowie Nahrungs- «FIII ID» 9.3 × Nahrungsergänzungs- und«/RII» Feed, which < at least a connection of the general formula; RTI ID» 9.4 > (I) < / >
«/RTI» contained.

A further subject matter of the invention is therefore a method < to the production of the inventive bonds of the formule; RTII De-95.5 (I) < ATIN by the fact the characterized is that a sugar 2 and/or, a Glykodid < RTI ID-9.65 Z-O-Pm</ ATI) mit einer ungesättigten Fattsäure ARTI ID-9.75 Ac0Hz/RTI> oder mit einem Ester, vorzugsweise einem Methy-loder Birbylester, dieser Fettsäuren Acolf in Gegenwart einer Upsas versetert wird.

on the suitable arraymatic catalysts to the esterification (by means of transesterification) the hydroizes, the particular Lipseer make of the acids and alcohol components mentioned like the Lipseer from Candida exterior. As a particular Lipseer from Candida cyloradida rugosa (Index) (and Lipseer from Candida cyloradida rugosa (Index) (and Lipseer from Candida cyloradida rugosa (Index) (and Lipseer from Candida cyloradida rugosa) (Index) (Ind

A preferred Upase is the Lipase (ISO enzyme B) from Candida antarctica, for which there are two reasons. First of all it particularly shows one high selectivity with the esterification of the acetals with the insatiated tatty acids, although these do not rank among their typical substrates. Des weiteren zeigt sie keine Grenzflächenaktivierung (ein entscheidendes Merkmal zur Klassel-fizierung von Hydrolasen in die Gruppe der Lipasen), da hir ein wichtiges «RII ID=9.9» Lipasestrukturmerk-mal.</br>
/RII > 9.9 » Lipasestrukturmerk-mal.

Es wird also die selektive Veresterungseigenschaften einiger Enzyme (siehe

Beschreibung), insbesondere Lipase aus Candida < RTII ID=10.15 antarctica Iscenzym B,</RTI> in addition used,
To link and partly improve active substances with one another to esters and so the effects and a simultaneous possibility
to have of isolating the most active isomer of the CLAs from the mixture.

Gerade die gezielte Veresterung ist aber für die biologische Verfügbarkeit und

Compatibility of the inventive materials crucial. The chemical

Synthese führt jedoch aufgrund mangelnder Regioselektivität zu groben Poduct mixtures. Therefore is the here described enzymatic (see examples) mild and regioselektive synthesis of advantage. Inventive one means regiospezilisch that only a certain OH-group of a polyole becomes third.

Corresponding one means regioselektiv that a certain OH-group of a polyole does not become prefered, but exclusive,

The inventive connections are < RTI ID=10.2> Formula (I) < /RTI> erts einmal mittels doe erindungsgemäsen Verlahrens hergestellt worden, muss in aller Regel ein Verlahren folgen, <RTI ID=10.3> un=6 leg ewünsche</RTI> (n) To up-to-some connection (EN). Thus a further subject matter of the present invention consists of it, a method to the purification of the bonds of the formula < RTI ID=10.4> (I) < /RTII> to make available, which is characterized thereby that it concerns an aqueous two-phase extraction procedure with organic solvents, with which the target compound from the not converted falty acids separated become selective can. Vorzugeweise handelt es ein bei dem organischen Lösungsmittel um n-Hexan. Cyclohexan, THF, Dieethylether. Alternative one knows the purification also by a chromatographic method at silicagel, preferably with < RTI ID=10.5> Ethyl scateate/methanol order: /RTI> Dichloromethane/methanol mixtures with minor proportions acetic acid and/or water takes place, which can become also additional an aqueous two-phase extraction procedure with organic solvents performed.

Since the inventive esters of the formula < RTI ID=10.6> (I) < /RTI> a good biological availability and effect have, can it in cosmetic and pharmaceutical preparing and/or as food additives be used with the result that the quality of these products become remarkably improved.

Die erfindungsgemässen Ester der konjugierten Linolsäuren (CLAs) weisen eine «RTI ID=11.1> antlimitiamerfinden, erfinden, erfindungsgemässen Ester der konjugierten angeleische Wirkung auf und haben über-dies auch antioxidative, «RTI ID=11.2> hautaulfhellende-(RTI) or antibacterial/artiviral effects. Die «RTI ID=11.3> Fettdepotreduzierende-(RTI). Effect of the CLAs shows until inventive connection.

Since the connections of the formula $< R\Pi | D=11.4> (I) < /R\Pi >$ a good biological availability and effect have, can it in cosmetic and pharmaceutical preparing and/or as food additives be used with the result that the quality of these products becomes remarkably improved.

Beyond that the inventive connections are particularly good into lipophilic

Basis prescriptions trainable and can be formulated in a simple manner as stable emulsions.

Accordingly the inventive connections of the formula are < RTI ID=11.5> (I) < /RTI> to the production of cosmetic and/or pharmaceutical preparing and/or Nahrungs-bzw. Feeds uses.

Further subject matters of the invention are < thereafter the use of the connections; RII ID=11.6.5 Formula (I) < /RII IT and IT is a first threat of the connections; RII ID=11.6.5 Formula (I) < /RII IT is a first threat of the connections; RII ID=11.6.5 Formula (II) < RII IT is a first threat of the connection (II) of the first threat of the connection (II) of the formula; RII III D=11.7.5 (I) < RIII > contained.

Die unter erfindungsgemässer Verwendung der Verbindungen «RII ID=11.8»(I)
 /RII) a variable cosmetic preparing such as hair shampoos, «RII ID=11.9» Hair lotions, «/RII) = Foam baths, «RII ID=11.0» Liber lotions, «/RII) = Foam baths, «RII ID=11.10» (RII) = Foam baths, «RII ID=11.10» (RII) = Foam baths, «RII ID=11.11» (RII) = Foam baths, «RIII) = Foam baths, «RIII = Foam baths, »RIII = Foam baths, «RIII = Foam baths, «RIII = Foam baths, «RIII = Foam baths, »RIII = Fo

Hydrotrope, < RTI ID=12.1> Preservative, Insektenrepellentien, selfbrown, Solubilisatoren, < /RTI> Perfume oils, dyes, germ-inhibitive means and such contained.

The quantity required of the inventive connections in the cosmetic preparing lies usually within the range of 0,01 to 5 Gew. - %, preferably however from 0,1 to 1 Gew. - % related to the total quantity of the preparing.

The daily dosage required for the achievement of a corresponding effect with pharmaceutical applications appropriately 0.1 to 10 amounts to mg/kg body weight, preferably 0.5 to 2 mg/kg body weight.

The food spare and additives as, available under inventive use of the connections of the formula (I), < RTI ID=12.5> Sportier Drinks</ RTI> contained suitable-proves the connection (RN) < RTI ID=12.6> Formula (I), < RTII> in a quantity, which leads with a usual need at Rüssigkietsaufnahme from 1 to 5 iltres per day to a dosage of these bonds from at 0.1 to 10 mg, preferably 0.5 to 5 mg, per kg body weight. An exemplary use in food industry exists for the connections of the formula < RTII D=1.27.9 (I) < RTII> < RTII D=1.28> as Fārbe-undfor Gewürzstelfie. < RTII>

Examples example 1: CLA+ glucose < RTI ID=13.15 6-O-cis9, < /RTI s trans-11-Octadecadiencyi-D-glucopyranose 136 (0.1 mol) < RTI ID=13.25 D (+): glucose, < /RTI > 39.6 g CLA (60% conjugated Linoisäure) (Edenor UKD 6010), 30 < RTI ID=13.35 g < /RTI > Molecular sieve, 120 ml I: Butanol and 3g immobilized Upase B out Candida antartica were < 48 hours with 60 C with 400 RPM at the magnet agitator in 300; RTI ID=13.45 ml Efrenmever/kolber / RTI is inkubiert. The conversion became by means of

Thin layer chromatography (silicagel 80-Platten with < RTI ID=13.5: Pluoreszenzindikator < RTI > 1.2 utilities (< RTI > 1.3 utilities (<

RI-value: 0.41 c. RTI ID=13.9c. (Ehyl acetate/Methanolc. /RTI> 10: 1) 0.05 (diolhoromethan/methanol/acetia cide 57: 5: 0.1) c. RTI ID=13.19c. 13G-NMRc /RTI> (100.6 MHz, CD30D); c. RTI ID=13.11> 8c./RTI> 8c./RTI

93.8 (C-1), 129.7 < RTI (ID=13.21> (C-10), < / RTI> 130.2 < ŘTI (ID=13.22> (C-11), < / RTI> 131.6 (C-9), 173.6 (c1).

< RTI (ID=13.24> 8c. / RTI = (6is cis isomen) 14.4 (C-18), 23.8 (C-17), 25.6 (C-3) 20.1-30.7 (C-4, C-5, C-6, C-7, <- RTI (ID=13.24> 6c./ RTI) = (6is cis isomen) 14.4 (C-18), 24.8 (C-17), 25.6 (C-3) 20.1-30.7 (C-4, C-5, C-6, C-7, <- RTI (ID=13.26> (C-13), <- RTI) = 33.6 (C-6), 34.4 (C-2), 64.8 (C-7).

ID=13.27> (C-6'), < /RTI> 70.4 < RTI ID=13.20> (C-4'), < /RTI> 71.8 (C< RTI ID=13.20> 5'), < /RTI> 73.8 < RTI ID=13.20> (C-2'), < /RTI> 74.9 < RTI ID=13.31> (C-2'), < /RTI> 74.9 < RTI ID=13.31> (C-2'), < /RTI> 74.8 (cf).

 $RI-value: 0.74 < RTI \; ID=14.3> \; (Ethyl \; acetate/Methanol < /RTI> \; 10:1) < RTI \; ID=14.4> \; Example \; 3: \\ Stear idons \\ aureethylester < /RTI> + \; Arbutin \; Arb$

69, 12,15-Octadecatetraenoyl (62, 92, < RTI ID=14.5> 122, 152) - Arbutin</ri>
(7RTI> 1,36 g (5 mmol) Arbutin, < RTI ID=14.6> 40g Stearidonsäureethylester, < /RTI> 4.5 g molecular sieve, 10 ml 1-Butanol and 4,5 g immobilized Lipase B from Candida antartica were < 48 hours with 60; RTI ID=14.7> C> (RTI) with 100 RPM at the magnet agitator in 300 ml Erlomneyer (lasks inkubiert. The conversion became by means of thin layer chromatography (silicagel < RTI ID=14.8> 60 PAttern < /RTI> but filtorescence indicator; Lautimitte: Ethylacetat 100 %, < RTI ID=14.8> 60 PAttern < /RTI> but filtorescence indicator; Lautimitte: Ethylacetat 100 %, < RTI ID=14.8> 60 PAttern < /RTI> but filtorescence indicator; Lautimitte: Ethylacetat 100 %, < RTI ID=14.8> 60 PAttern < RTIS PATTERN </RTIS PATTERN </RTIS PATTERN </RES PATTERN </ri>

| ID=14.9>Visualisierung</RIT>: UV detection as well as by means of acetic acid/sulphuric acid/anisic aldehyde dipping reagent (100: 2: 1, < RIT ID=14.10> v/v/v)) / /RIT> proven. The product became over Säulenchroma-tographie (silicage) #676; Ludr/HITE IIII publicedti2-Poponal 10: 1, v/v) purified.

<RTI ID=14.11>Ru-vert</RTI>: 0.35 (ethyl acetate 100%)



Claims of W 00179241 Print Copy Contact Us Close

Result Page

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Claims

- Connections of the general formula (1): < RTI ID=15.1> Acr, O-Z-0-R, < /RTI> (1), where R a hydrogen, a one < over one; RTI ID=15.2> Etherbrückee / RTI> at the sugar bound branched or straight-chain < RTI ID=15.3> Go-Zo2-Alkki. < /RTI> one Arvialkyl or a substituded or a unsubstitulerten < RTI ID=15.4> Go-C. o-arly radical represents.
- < /RID: where Z (sugar) stands for a mono, a disaccharide or a polysaccharide, which are n-multiple ester-like with AC usubstituted and, if R does not stand for hydrogen, acetalisch to the remainder R bound, where AC a multiple installated < RTI ID-15.5> C5-C25-Acylrest
 /RTI ID-15.5> C5-C25-Acylrest
 /RTI D-15.5> C5-C25-Acylrest
 /RTI D-15.6> C5-C25-Acylrest
 /RTI D-15.6
 /RTI D-15.
- 2. The bonds of claim 1, whereby Z a monosaccharide, in particular D
- Glucose, D-Galactose, D-xylose, D-Apiose, L-Rhamnose, L-arabinose and Rutinose, are.
- 3. The connections of claim 1 or 2, whereby < RTI ID=15.7> Glykosid portion (Z-O-Rm) < /RTI> such is, that as a sugar component a monosaccharide, in particular in Monosaccharide according to claim 2, contained bonds, in particular
- 4. The connections of claim 1 or 2, whereby the Glykosid portion < RTI ID=15.9> (Z-O-Rm) < /RTI> such is, that as a sugar component Oligo or a polysaccharide how Alkylpolyglucoside, particular APG < RTI ID=15.10> (Handle KGaA), < /RTI> is.
- 5. The bonds of one of the preceding claims, whereby the linkage of AC at the sugar over a primary alcohol group of the sugar made.
- 6. The connections of one of the preceding claims, whereby that Glykosid

Arbutin, < RTI ID=15.8> Fragilin< /RTI> and Poplin.

- Portion < RTI ID=16.1> (Z-O-Rm) < /RTI> Arbutin, Populin or < RTI ID=16.2> Fragilin< /RTI> is.
- 7. The connections of one of the preceding claims, whereby AC is the acyl radical of one of the subsequent fatty acids: a CLA or a Stearidonsaure, in particular those
- Octadecadiensäuren, those over conjugated double bonds at the C-atoms 9 and
- 11,10 and 12 or 11 and 13 orders, in particular cis, trans, 9,11-und cis, cis, 9,11-Octadecadiensaure.
- 8. The connections of one of the preceding claims, whereby that Glykosid
- Portion < RTI ID=16.3> (Z-O-Rm) < /RTI> Arbutin (D. h. m = 1), whereby AC the acyl radical cis, trans, 9,11-oder cis, cis, 9,11-Octadecadiensaure and whereby n 1 is.
- 9. The connections of one of the claims 1 to 7, whereby the Glykosid portion < RTI ID=16.4> (Z-O-< /RTI> \$\frac{10p}{3} Rm) Fragulin (D. h. m = 1), whereby AC the acyl radical cis, trans, 9,11-oder cis, cis,
- 9,11-Octadecadiensaure and whereby n 1 is.
 - < RTI ID=16.5> 10.
 /RTI> The connections of one of the claims 1 to 7, whereby < RTI ID=16.6> Glykosid portion (Z-0/RTI>
 - Rm) Arbutin (D. < RTI ID=16.7> h.< /RTI> m = 1), whereby AC the acyl radical cis, trans, 9,11-oder cis, cis, 9,11-Octadecadiensäure and whereby n 2 is.
 - < RTI ID=16.8> 11.< /RTI> The connections of one of the claims 1 to 7, whereby the Glykosid portion < RTI ID=16.9> (Z-O-</RTI>
 - Rm) Fragulin (D. h. m = 1), whereby AC the acyl radical cis, trans, 9,11-oder cis, cis,
 - 9,11-Octadecadiensäure and whereby n 2 is.
 - 12. The connections of one of the claims 1 to 7, whereby < RTI ID=16.10> Glykosid portion (Z-O-< /RTI>
 - Rm) Arbutin or Fragilin (D. h. m = 1), whereby AC the acyl radical of the Stearidonsaure and whereby n 1 is.
 - RTI ID=17.15 13.
 /RTI> The connections of one of the claims 1 to 7, whereby the Glykosid portion (Z-O Rm) Arbutin or
 RTI ID=17.2> Fragulin
 /RTI> (D. h. m = 1), whereby AC the acyl radical of the Stearidonsäure and whereby n 2 is.
 - 14. Connections of one preceding claims, whereby, if n = 1, which is single remainder of AC to a primary OH-group of the sugar bound, in particular to that primary OH-group, which sits at the sugar moiety, to which also R bound it is, if m < RIT ID=17.3> w< /RIT> 0, and whereby, if n = 2, a remainder of AC to that primary OH-group, which sits at the sugar moiety, to which also R bound is, if m < RIT ID=17.4.5 # < /RIT> 0, and which is other remainder of AC to a further primary or to a secondary OH-group of the sugar bound.

- $< RTI\ ID = 17.5 >\ 15. </RTI> \ Method to the production of the bonds < RTI\ ID = 17.6 >\ Formula\ (I) </RTI> of one of the preceding claims, characterised in that a sugar Z and/or.$
- Glykosid < RTI ID=17.7> Z-O-Rm< /RTI> with an insatiated fatty acid < RTI ID=17.8> AcOH< /RTI> or with an ester, preferably one methyl or ethyl ester, these fatty acids < RTI ID=17.9> AcOH< /RTI> in presence of a hydrolase third or one unnecester.
- 16. The method of claim 15, whereby the hydrolase a Lipase, in particular a Lipase from Candida rugosa (formerly Candida cylindaceae), Candida antarctica, Geotrichum candidum, aspergillus niger, Penicillium roqueforti, Rhizopus arrhizus and Mucor miehei, in particular the Lipase < RII 105 17.10> (Isoportym < IRIN) = By from Candida antarctica, is.</p>
- 17. The method of claim 15 or 16, whereby itself to the esterification reaction
- Step to the purification of the bonds of the formula (I) attaches, either the aqueous two-phase extraction procedure with organic solvents such as n-hexane, eydohoxano, HF or < RII | 10 = 17.11.0 | bethylether < |RII| > or a chromatographic Method at silicagel, preferably with < RII | 10 = 17.12.0 | Ethyl acotate/methanol oder < / | /RII| > or a chromatographic Dichloromethane/methanol mixtures with minor proportions accelic acid and/or
- 18. Cosmetic or pharmaceutical composition or Nahrungs-bzw.

Nahrungsergänzungs or feed composition, contained at least one of the connections of one of the claims 1 to 14.